ANSWER 1 OF 4 CA COPYRIGHT 2003 ACS 137:334779 CA AN FAST CARS: Engineering a laser spectroscopic technique for rapid identification of bacterial spores Scully, M. O.; Kattawar, G. W.; Lucht, R. P.; Opatrny, T.; Pilloff, H.; ΑU Rebane, A.; Sokolov, A. V.; Zubairy, M. S. Institute for Quantum Studies, Texas A and M University, College Station, CS TX, 77843, USA Proceedings of the National Academy of Sciences of the United States of SO America (2002), 99(17), 10994-11001 CODEN: PNASA6; ISSN: 0027-8424 National Academy of Sciences PB Journal DT English LA Airborne contaminants, e.g., bacterial spores, are usually AB analyzed by time-consuming microscopic, chem., and biol. assays. Current research into real-time laser spectroscopic detectors of such contaminants is based on e.g., resonance fluorescence. The present approach derives from recent expts. in which atoms and mols. are prepd. by one (or more) coherent laser(s) and probed by another set of lasers. However, generating and using maximally coherent oscillation in macromols. having an enormous no. of degrees of freedom is challenging. In particular, the short dephasing times and rapid internal conversion rates are major obstacles. However, adiabatic fast passage techniques and the ability to generate combs of phase-coherent femtosecond pulses provide tools for the generation and utilization of maximal quantum coherence in large mols. and biopolymers. We call this technique FAST CARS (femtosecond adaptive spectroscopic techniques for coherent anti-Stokes Raman spectroscopy), and the present article proposes and analyses ways in which it could be used to rapidly identify preselected mols. in real time. CC 9-5 (Biochemical Methods) FAST CARS engineering laser spectroscopic technique bacterial ST spore Bacillus cereus IT Bacillus megaterium Bacteria (Eubacteria) CARS Raman spectroscopy Laser spectroscopy Resonance fluorescence Spore (FAST CARS for rapid identification of bacterial spores) TT Biopolymers Macromolecular compounds RL: ANT (Analyte); ANST (Analytical study) (FAST CAPS for rapid identification of bacterial spores) IT RL: BSU (Biological study, unclassified); BIOL (Biological study) (FAST CARS for rapid identification of bacterial spores) Proteins PL: BSU (Biological study, unclassified); BIOL (Biological study) (FAST CARS for rapid identification of bacterial spores) IT PL: BSU (Biological study, unclassified); BIOL (Biological study) (FAST CARS for rapid identification of bacterial spores) ΙT 6893-30-7, Calcium dipicolinate RL: ARU (Analytical role, unclassified); ANST (Analytical study) (FAST CARS for rapid identification of bacterial spores) RE.CNT THERE ARE 66 CITED REFERENCES AVAILABLE FOR THIS RECORD 66

L11 ANSWER 2 OF 4 CA COPYRIGHT 2003 ACS

AN 134:233815 CA

TI Physical perturbation for fluorescent characterization of microorganism

ALL CITATIONS AVAILABLE IN THE RE FORMAT

particles Bronk, Burt V.; Shoaibi, Azadeh; Nudelman, Raphael; Akinyemi, Agnes N. ΑU AFRL/ECBC at U.S. Army ECBC, A.P.G., MD, 21010-5424, USA CS Proceedings of SPIE-The International Society for Optical Engineering SO (2000), 4036 (Chemical and Biological Sensing), 169-180 CODEN: PSISDG; ISSN: 0277-786X PВ SPIE-The International Society for Optical Engineering DT Journal English LA AB The motivation for using response to phys. perturbation to classify microparticles came from our previous expts. with Dipicolinic Acid (DPA). DPA as a calcium complex is a major component of bacterial spores, constituting more than 5% of their dry wt. It is not commonly found in other natural products and therefore its presence is indicative of the presence of bacterial spores. Previous schemes utilizing the presence of DPA to detect these spores have relied on **fluorescence** which occurs when lanthanide metals (e.g., terbium) are added to a soln. where the presence of DPA is to be detd. We have recently demonstrated that changes in the fluorescence of DPA can be stimulated without the addn. of such reagents. Thus after exposure to UV light, a substantial increase of fluorescence emitted by DPA solns. with a peak at 410 nm occurs for excitation light with wavelength less than approx. 305 nm. 9-5 (Biochemical Methods) Section cross-reference(s): 10 fluorometry microorganism optical classification dipicolinate ST Bacillus megaterium ΙT Bacillus subtilis Escherichia coli Fluorometry Microorganism (phys. perturbation for fluorescent characterization of microorganism particles) TT 499-83-2, Dipicolinic Acid RL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses) (phys. perturbation for fluorescent characterization of microorganism particles) THERE ARE 11 CITED REFERENCES AVAILABLE FOR THIS RECORD RE.CNT 11 ALL CITATIONS AVAILABLE IN THE RE FORMAT L11 ANSWER 3 OF 4 CA COPYRIGHT 2003 ACS AN132:75561 CA Dipicolinic acid (DPA) assay revisited and appraised for ΤI spore detection AII Hindle, Alistair A.; Hall, Elizabeth A. H. Inst. Biotechnol., University of Cambridge, Cambridge, CB2 1QT, UK CS Analyst (Cambridge, United Kingdom) (1999), 124(11), 1599-1604 SO CODEN: ANALAO; ISSN: 0003-2654 PΒ Royal Society of Chemistry DT Journal LA English Delayed gate fluorescence detection of dipicolinic AΒ acid (DPA), a universal and specific component of bacterial spores , has been appraised for use in a rapid anal. method for the detection of low concns. of bacterial spores. DPA was assayed by fluorimetric detection of its chelates with lanthanide metals. The influence of the choice and concn. of lanthanide and buffer ions on the fluorescence assay was studied as well as the effects of pH and temp. The optimal system quantified the fluorescence of terbium monodipicolinate in a soln. of 10  $\cdot$ mu.M terbium chloride buffered with 1 M sodium acetate, pH 5.6 and had a detection limit of 2 nM DPA. This assay allowed the first real-time monitoring of the germination of bacterial spores by continuously quantifying exuded DPA. A detection limit

```
of 104 Bacillus subtilis spores ml-1 was reached, representing a
     substantial improvement over previous rapid tests.
     9-5 (Biochemical Methods)
CC
     Section cross-reference(s): 10
     dipicolinate DPA detn Bacillus spore fluorometry
ST
ΙT
     Bacillus subtilis
     Fluorometry
     Solvent effect
       Spore
        (dipicolinic acid (DPA) assay for spore detection)
     499-83-2, Dipicolinic acid
     RL: ANT (Analyte); BOC (Biological occurrence); BSU (Biological study,
     unclassified); ANST (Analytical study); BIOL (Biological study);
     OCCU (Occurrence)
        (dipicolinic acid (DPA) assay for spore detection)
     13759-92-7, Europium chloride hexahydrate 13798-24-8, Terbium chloride
TT
     hexahydrate 15059-52-6, Dysprosium chloride hexahydrate
     RL: ARG (Analytical reagent use); ANST (Analytical study); USES
     (Uses)
        (dipicolinic acid (DPA) assay for spore detection)
              THERE ARE 51 CITED REFERENCES AVAILABLE FOR THIS RECORD
        51
              ALL CITATIONS AVAILABLE IN THE RE FORMAT
    ANSWER 4 OF 4 CA COPYRIGHT 2003 ACS
L11
     131:141627 CA
ΤI
     Fluorescence of dipicolinic acid as a possible
     component of the observed UV emission spectra of bacterial spores
     Nudelman, Raphael; Feay, Nicole; Hirsch, Mathew; Efrima, Shlomo; Bronk,
AU
    Mantech Environmental Technology Inc., USA
CS
     Proceedings of SPIE-The International Society for Optical Engineering
SO
     (1999), 3533(Air Monitoring and Detection of Chemical and Biological
    Agents), 190-195
    CODEN: PSISDG; ISSN: 0277-786X
    SPIE-The International Society for Optical Engineering
PΒ
DT
    Journal
LA
    English
    Dipicolinic acid (DPA) and the Ca2+ complex of DPA (CaDPA) are
     well-known and are major chem. components of bacterial spores.
     DPA's native fluorescence is very weak and is thought to be
     completely masked by the fluorescence of tryptophan when this
     compd. is present. Thus fluorescence related to DPA in
     spores is assumed by many authors to be completely absent. We
     show that the fluorescence of CaDPA is substantial for
    excitation between about 290 nm and 310 nm with emission peaking near 400
    nm. This emission is at the long wavelength tail for emission from
    tryptophan. We examine whether the emission of CaDPA could contribute to
    the total emission spectrum when bacterial spores are present in
    an aerosol, for excitation wavelengths in the neighborhood of 310 nm.
    this report we present measurements of fluorescence excitation
    and emission for CaDPA and compare them with that of DPA and tryptophan.
    9-5 (Biochemical Methods)
    Section cross-reference(s): 4, 10
ST
    bacterium spore dipicolinate fluorometry
    Bacteria (Eubacteria)
    Environmental analysis
    Fluorometry
      Spore
        (fluorescence of dipicolinic acid as a possible
       component of obsd. UV emission spectra of bacterial spores)
    73-22-3, L-Tryptophan, analysis 499-83-2, Dipicolinic acid
IT
    499-83-2D, Dipicolinic acid, calcium complex
                                                   7440-70-2D,
    Calcium, complex with dipicolinic acid, analysis
    RL: ANT (Analyte); BOC (Biological occurrence); BSU (Biological study,
```

unclassified; ANST (Analytical study); BIOL (Biological study);
OCCU (Occurrence)

(fluorescence of dipicolinic acid as a possible component of obsd. UV emission spectra of bacterial spores:

RE.CNT 15 THERE ARE 15 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

```
L19 ANSWER 4 OF 4 CA COPYRIGHT 2003 ACS
AN
     123:328802 CA
    Native fluorescence detection and spectral differentiation of
TI
     peptides containing tryptophan and tyrosine in capillary electrophoresis
AU
     Timperman, Aaron T.; Oldenburg, Kurt E.; Sweedler, Jonathan V.
     Department of Chemistry, University of Illinois, Urbana, IL, 61801, USA
CS
     Analytical Chemistry (1995), 67(19), 3421-6 CODEN: ANCHAM; ISSN: 0003-2700
SO
PΒ
     American Chemical Society
DT
     Journal
LA
    English
     A native fluorescence detection system for capillary
AB
     electrophoresis is described that achieves low attomole detection limits
     and simultaneous acquisition of complete fluorescence
     emission spectra. The system is designed for detection of
     peptides through the intrinsic fluorescence of
     tryptophan and tyrosine residues. The detection system employs a
     frequency doubled krypton laser operating at 284 nm for excitation
     , a sheath flow cell, a reflective f/1.2 microscope objective, an imaging
     spectrograph, and a CCD detector. The detection capabilities were
     characterized with tryptophan and tyrosine, which have limits of detection
     (3.sigma.) of 2 .times. 10-10 and, 2 .times. 10-8 M, resp. Acquisition of
     the fluorescence emission spectrum provides the
     ability to distinguish three classes of peptides: those that contain
     tryptophan, those that contain tyrosine, and those that contain both
     tryptophan and tyrosine.
     80-2 (Organic Analytical Chemistry)
CC
     Section cross-reference(s): 9
     native fluorescence detection tryptophan tyrosine peptide;
ST
     electrophoresis fluorescence detection tryptophan tyrosine
     peptide
     Fluorometers
TT
        (for native fluorescence detection and spectral
        differentiation of peptides contg. tryptophan and tyrosine in capillary
        electrophoresis)
ΙT
     Peptides, analysis
     RL: ANT (Analyte); ANST (Analytical study)
        (native fluorescence detection and spectral differentiation
        of peptides contq. tryptophan and tyrosine in capillary
        electrophoresis)
ΙT
     Electrophoresis and Ionophoresis
        (capillary, detectors, for native fluorescence detection and
        spectral differentiation of peptides contg. tryptophan and tyrosine)
    Microbial hormones and pheromones
IT
     RL: ANT (Analyte); ANST (Analytical study)
        (.alpha.-factor, native fluorescence detection and spectral
        differentiation of peptides contg. tryptophan and tyrosine in capillary
        electrophoresis)
     60-18-4D, Tyrosine, peptides
                                   73-22-3, Tryptophan, analysis
TT
                           21778-69-8
                                         58822-25-6, Leucine-enkephalin
     Tryptophan, peptides
     59401-28-4, .alpha.-Mating factor (yeast) 65418-88-4, .alpha.-1-Mating
              89911-64-8, Cholecystokinin(26-31)
                                                   98395-75-6, Neuromedin U 8
     (pig spinal cord)
     RL: ANT (Analyte); ANST (Analytical study)
        (native fluorescence detection and spectral differentiation
        of peptides contg. tryptophan and tyrosine in capillary
```

electrophoresis)

```
L17
    ANSWER 5 OF 179 CA COPYRIGHT 2003 ACS
    138:85885 CA
AN
    Comparative intrinsic and enhanced total photoluminescence of endospore
ΤI
    material
     Anderson, John E.; Webb, Stanley Thomas; Fischer, Robert L.; Kester,
ΑU
     Karen; Smith, Clint
     U.S. Army Topographic Engineering Center, Alexandria, VA, 22315, USA
CS
     Proceedings of SPIE-The International Society for Optical Engineering
SO
     (2002), 4576 (Advanced Environmental Sensing Technology II), 27-31
     CODEN: PSISDG; ISSN: 0277-786X
    SPIE-The International Society for Optical Engineering
PB
DT
    Journal
    English
LA
    Two techniques are compared using total luminescence spectroscopy to
AB
     detect endospore material in prepns. equiv. to 3.0 x 105/mL spores
       The first method applied intrinsic, steady-state photoluminescence for
     detection. The second approach using a binding fluorochrome derived from
     4-p-dimethylaminostyrylpyridinium (DASP) to signal the presence of
     spore material. Comparative fluorescence
     emission signatures (excited at 469 nm) showed greater calibrated
     signal recovery (4x106 cps) for spore material at longer
     wavelengths using DASP. The intrinsic fluorescence
     emission of endospores (excited at 346 nm) occurred at shorter
     wavelengths and showed a reduced calibrated intensity (1.4 x 105 counts
     per s, cps). One major advantage of DASP appears to be its longer
     wavelength excitation (469 nm) that is out of the range of
     assocd. biol. materials that compete for absorption at shorter UV
    wavelengths.
    9-5 (Biochemical Methods)
CC
    photoluminescence spectroscopy endospore
ST
ΙT
    Luminescence spectroscopy
        (comparative intrinsic and enhanced total photoluminescence of
       endospore material using dimethylaminostyrylpyridinium fluorochrome)
ΙT
     Fluorescent dyes
        (dimethylaminostyrylpyridinium deriv.; comparative intrinsic and
        enhanced total photoluminescence of endospore material using
       dimethylaminostyrylpyridinium fluorochrome)
IT
        (endospore; comparative intrinsic and enhanced total photoluminescence
       of endospore material using dimethylaminostyrylpyridinium fluorochrome)
TT
     186659-60-9
     RL: ARG (Analytical reagent use); ANST (Analytical study); USES
     (Uses)
        (fluorochrome derived from; comparative intrinsic and enhanced total
       photoluminescence of endospore material using
       dimethylaminostyrylpyridinium fluorochrome)
             THERE ARE 10 CITED REFERENCES AVAILABLE FOR THIS RECORD
             ALL CITATIONS AVAILABLE IN THE RE FORMAT
L17 ANSWER 12 OF 179 CA COPYRIGHT 2003 ACS
AN
    137:286140 CA
ΤI
    Optical structure for multi-photon excitation and the use
    thereof
ΤN
    Duveneck, Gert L.; Bopp, Martin A.; Pawlak, Michael; Ehrat, Markus;
    Marowsky, Gerd
    Zeptosens A.-G., Switz.
PΑ
SO
    PCT Int. Appl., 76 pp.
    CODEN: PIXXD2
DT
    Patent
    German
LA
FAN.CNT 1
                           DATE APPLICATION NO. DATE
    PATENT NO. KIND DATE
```

```
A2
    WO 2002079765
                           20021010
                                          WO 2002-EP2958
                                                           20020318
PΙ
                           20030130
    WO 2002079765
                     A3
        CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR,
            BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
PRAI CH 2001-617
                     Α
                           20010402
                           20010412
    CH 2001-689
                      Α
    Optical structures comprising an optical waveguide with a waveguiding
AB
    layer that is transparent at .gtoreq.l excitation wavelength are
    described in which the intensity of excitation light that is
    input into and guided via the waveguiding layer is sufficiently high to
    excite mols. or groups of mols. at a surface of the layer or within 200 nm
    of the layer by means of multi-photon excitation, preferably
    two-photon excitation. Optionally, an adhesive agent may be
    used to immobilize biochem. or biol. mols., or other materials, for use in
    detecting or detg. analytes in a sample. Optical systems for carrying out
    multiphoton excitation, and methods for anal. (e.g.,
    luminescence excitation and to the luminescent detection of one
    or several analytes) using the systems are also described. The systems
    may also be used to form optical tweezers. Methods for luminescence and
    fluorescence anal. of biomols., including autofluorescence of
    nucleic acids, is emphasized.
TC
    ICM G01N021-77
    ICS G01N021-55; G01N021-64
    73-10 (Optical, Electron, and Mass Spectroscopy and Other Related
CC
    Properties)
    Section cross-reference(s): 9, 17, 64, 74
    waveguide multiphoton excitation system; luminescence analysis
ST
    waveguide multiphoton excitation system; fluorescence
    analysis waveguide multiphoton excitation system
    Prion proteins
IT
    RL: ANT (Analyte); ANST (Analytical study)
        (PrPSc, assay for; wavequide structures for multiphoton
       excitation and methods of multiphoton excitation
       using them and use of the structures in luminescent anal.)
ΙT
    Bacteria (Eubacteria)
    Pathogen
    Salmonella
        (assay for; waveguide structures for multiphoton excitation
       and methods of multiphoton excitation using them and use of
       the structures in luminescent anal.)
TΤ
    Fluorescence
        (autofluorescence; wavequide structures for multiphoton
       excitation and methods of multiphoton excitation
       using them and use of the structures in luminescent anal.)
ΙT
    Disease, plant
        (diagnosis of; waveguide structures for multiphoton excitation
       and methods of multiphoton excitation using them and use of
       the structures in luminescent anal.)
ΙT
    Immunoassay
        (fluorescence; waveguide structures for multiphoton
       excitation and methods of multiphoton excitation
       using them and use of the structures in luminescent anal.)
TT
    Immunoassay
       (immunofluorometric; waveguide structures for multiphoton
       excitation and methods of multiphoton excitation
       using them and use of the structures in luminescent anal.)
ΙT
    Photoexcitation
```

(multiphoton; wavequide structures for multiphoton excitation and methods of multiphoton excitation using them and use of the structures in luminescent anal.) Polyesters, uses RL: DEV (Device component use); USES (Uses) (thio-; waveguide structures for multiphoton excitation and methods of multiphoton excitation using them and use of the structures in luminescent anal.) TT Photoexcitation (two-photon; waveguide structures for multiphoton excitation and methods of multiphoton excitation using them and use of the structures in luminescent anal.) Blood analysis IT Combinatorial chemistry Diagnosis Drug design Drug screening Egg yolk Fluorescence Fluorometers Fluorometry Food analysis Gene expression profiles Luminescence Optical waveguides Pharmaceutical analysis Plant analysis Surface waters Urine analysis (waveguide structures for multiphoton excitation and methods of multiphoton excitation using them and use of the structures in luminescent anal.) Agglutinins and Lectins ΙT Carbohydrates, analysis Enzymes, analysis Oligonucleotides P.NA Receptors PL: ANT (Analyte); ANST (Analytical study) (waveguide structures for multiphoton excitation and methods of multiphoton excitation using them and use of the structures in luminescent anal.) Acrylic polymers, uses TТ FL: DEV (Device component use); USES (Uses) (waveguide structures for multiphoton excitation and methods of multiphoton excitation using them and use of the structures in luminescent anal.) Polyamides, uses IT RL: DEV (Device component use); USES (Uses) (waveguide structures for multiphoton excitation and methods of multiphoton excitation using them and use of the structures in luminescent anal.) IT Polycarbonates, uses PL: DEV (Device component use); USES (Uses) (waveguide structures for multiphoton excitation and methods of multiphoton excitation using them and use of the structures in luminescent anal.) TT Polyesters, uses PL: DEV (Device component use); USES (Uses) (waveguide structures for multiphoton excitation and methods of multiphoton excitation using them and use of the structures in luminescent anal.) ΙT Polyimides, uses

(wavequide structures for multiphoton excitation and methods of multiphoton excitation using them and use of the structures in luminescent anal.) ΙT Polythiophenylenes RL: DEV (Device component use'; USES (Uses) (waveguide structures for multiphoton excitation and methods of multiphoton excitation using them and use of the structures in luminescent anal.) Polyurethanes, uses ΙT RL: DEV (Device component use); USES (Uses) (wavequide structures for multiphoton excitation and methods of multiphoton excitation using them and use of the structures in luminescent anal.) 71-00-1D, L-Histidine, oligomers IT RL: ANT (Analyte); ANST (Analytical study) (waveguide structures for multiphoton excitation and methods of multiphoton excitation using them and use of the structures in luminescent anal.) ΙT 1313-96-8, Niobium oxide 1314-13-2, Zinc oxide, uses 1314-23-4, Zirconium oxide, uses 1314-61-0, Tantalum oxide 9002-88-4, Polyethylene 9003-01-4, Polyacrylic acid 9003-07-0, Polypropylene 9003-53-6, Polystyrene 9011-14-7, Polymethylmethacrylate 12055-23-1, Hafnium oxide 13463-67-7, Titanium oxide, uses 25038-59-9, Polyethylene terephthalate, uses RL: DEV (Device component use); USES (Uses) (waveguide structures for multiphoton excitation and methods of multiphoton excitation using them and use of the structures in luminescent anal.) L17 ANSWER 17 OF 179 CA COPYRIGHT 2003 ACS AN 136:382505 CA Device for monitoring cells ТT Pitner, J. Bruce; Hemperly, John Jacob; Guarino, Richard D.; Wodnicka, ΤN Magdalena; Stitt, David T.; Burrell, Gregory J.; Foley, Timothy G., Jr.; Beaty, Patrick Shawn PΑ Becton, Dickinson and Company, USA U.S., 42 pp., Cont.-in-part of U.S. Ser. No. 715,557. SO CODEN: USXXAM DT Patent LA English FAN.CNT 3 PATENT NO. KIND DATE APPLICATION NO. DATE \_\_\_\_\_ US 6395506 B1 20020528 US 1999-342720 19990629 EP 509791 A1 19921021 EP 1992-303391 19920415 PΤ B1 19960703 EP 509791 R: DE, FR, GB, IT CA 2066329 AA 19921019 CA 1992-2066329 19920416
JP 05137596 A2 19930601 JP 1992-98368 19920418
JP 07073510 B4 19950809
US 2002192636 A1 20021219 US 2002-109475 20020328
US 2002155424 A1 20021024 US 2002-116777 20020404

PRAI US 1991-687359 B1 19910418 US 1993-25899 A2 19930303 US 1996-715557 A2 19960918 US 1999-342720 A2 19990629 US 2000-642504 A2 20000818 US 2001-966505 A2 20010928 The present invention relates to methods for detection and evaluation of AΒ metabolic activity of eukaryotic and/or prokaryotic cells based upon their ability to consume dissolved oxygen. The methods utilize a luminescence detection system which makes use of the sensitivity of the luminescent emission of certain compds. to the presence of oxygen, which

RL: DEV (Device component use; USES (Uses)

```
quenches (diminishes) the compd.'s luminescent emission in a
     concn. dependent manner. Respiring eukaryotic and/or prokaryotic cells
     will affect the oxygen concn. of a liq. medium in which they are immersed.
     Thus, this invention provides a convenient system to gather information on
     the presence, identification, quantification and cytotoxic activity of
     eukaryotic and/or prokaryotic cells by detg. their effect on the oxygen
     concn. of the media in which they are present.
     ICM C12Q001-18
NCL 435032000
CC
     9-1 (Biochemical Methods)
     Section cross-reference(s): 1, 4
ST
     device monitoring cell
ΙT
    Plates
        (Microtitration; device for monitoring cells)
     Analytical apparatus
IT
     Antibiotics
      Bacteria (Eubacteria)
     Biological materials
     Blood
     Blood serum
     Cell
     Cell proliferation
     Chemicals
     Coating materials
     Composition
     Concentration (condition)
     Culture media
     Cytotoxicity
     Drugs
     Escherichia coli
     Eukaryota
     Extracellular matrix
       Fluorescence quenching
     Impermeability
     Insecta
     Light
     Liquids
     Luminescence
     Luminescence quenching
     Luminescence spectroscopy
     Luminescent substances
     Mathematical methods
     Metabolism
     Microorganism
     Molecules
     Particles
     Permeability
     Prokaryote
     Pseudomonas aeruginosa
     Radiation
     Reducing agents
     Respiration, animal
     Respiration, microbial
     Sensors
     Solutes
     Wavelength
     Wetting
    Yeast
        (device for monitoring cells)
TΤ
    Toxins
    RL: ADV (Adverse effect, including toxicity); BIOL (Biological study)
       (device for monitoring cells)
TT
    Reagents
     RL: ARG (Analytical reagent use); ANST (Analytical study); USES
```

```
(device for monitoring cells)
     Plastics, analysis
     FL: ARU (Analytical role, unclassified); ANST (Analytical study)
        (device for monitoring cells)
     Pubber, analysis
     FL: ARU (Analytical role, unclassified); ANST (Analytical study)
        (device for monitoring cells)
     Silicone rubber, analysis
     RL: ARU (Analytical role, unclassified); ANST (Analytical study)
        (device for monitoring cells)
     Growth factors, animal
ΙT
     PL: BSU (Biological study, unclassified); BIOL (Biological study)
        (device for monitoring cells)
     Collagens, biological studies
IT
     FL: BUU (Biological use, unclassified); BIOL (Biological study); USES
     ·Uses)
        (device for monitoring cells)
     Entactin
TТ
     FL: BUU (Biological use, unclassified); BIOL (Biological study); USES
     (Uses)
        (device for monitoring cells)
IT
     Laminins
     PL: BUU (Biological use, unclassified); BIOL (Biological study); USES
        (device for monitoring cells)
     Proteoglycans, biological studies
IT
     PL: BUU (Biological use, unclassified); BIOL (Biological study); USES
        (heparitin sulfate-contg.; device for monitoring cells)
     Optical detectors
IΤ
        (luminescence; device for monitoring cells)
     Animal cell
TT
        (mammal; device for monitoring cells)
     Amino acids, biological studies
IT
     PL: BUU (Biological use, unclassified); BIOL (Biological study); USES
        (nonessential; device for monitoring cells)
ΙT
     Collagens, biological studies
     PL: BUU (Biological use, unclassified); BIOL (Biological study); USES
        (type IV; device for monitoring cells)
                                         15158-62-0D, Tris-2,2'-
     1499-10-1, 9,10-Diphenylanthracene
TТ
     bipyridylruthenium (II), salts 36309-88-3, Tris-4,7-diphenyl-1,10-
     phenanthroline ruthenium (II) chloride
                                             50525-27-4, Tris-2,2'-
     bipyridylruthenium (II) chloride hexahydrate.
                                                    63373-04-6D,
     Tris-4,7-diphenyl-1,10-phenanthroline ruthenium (II), salts
     RL: APG (Analytical reagent use); ANST (Analytical study); USES
        (device for monitoring cells)
     7631-86-9, Silica, analysis
     FL: APU (Analytical role, unclassified); ANST (Analytical study)
        (device for monitoring cells)
     59-05-2, Methotrexate
                            151-21-3, Sodium dodecyl sulfate, biological
             865-21-4, Vinblastine
                                      7757-83-7, Sodium Sulfite
     Oxygen, biological studies 26628-22-8, Sodium Azide 35607-66-0,
                55268-75-2, Cefuroxime 85721-33-1, Ciprofloxacin
     Cefoxitin
     FL: BSU (Biological study, unclassified); BIOL (Biological study)
        (device for monitoring cells)
    57-92-1, Streptomycin, biological studies 113-24-6, Sodium pyruvate
ΙT
                                                   119978-18-6, Matrigel
     1397-89-3, Fungizone
                           1406-05-9, Penicillin
     141907-41-7, Matrix metalloproteinase
    RL: BUU (Biological use, unclassified); BIOL (Biological study); USES
     (Uses)
```

(Uses)

(device for monitoring cells)

RE.CNT 9 THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

LIT ANSWER 18 OF 179 CA COPYRIGHT 2003 ACS

AN 136:337350 CA

TI Method for detecting living cells in medium and measuring pH of medium

IN Kawasaki, Yukishige; Tsuji, Takashi; Kurane, Ryuichiro PA Sangyo Gijutsu Sogo Kenkyusho, Japan; Bioindustry Association; Mitsubishi

Chemical Corp.

SO Jpn. Kokai Tokkyo Koho, 12 pp.

CODEN: JKXXAF

DT Patent

LA Japanese

FAN.CNT 1

PATENT NO. KIND DATE APPLICATION NO. DATE

PI JP 2002125696 AZ 20020508 JP 2000-319247 20001019

PRAI JP 2000-319247 20001019

AB A convenient method is provided for simultaneously performing the detection of living cells (e.g, microorganism) in a medium and the measurement of the medium pH. The method comprises a step for adding a fluorescent enzyme substrate (e.g., 5-carboxyfluoroscein diacetate acetoxymethyl ester, 5-(and 6-)carboxyfluoroscein diacetate, 2',7'-bis-(2-carboxyethyl)-5-(and 6-)carboxyfluorescein acetoxymethyl ester, 5-sulfofluorescein diacetate) to the medium contg. living cells, a step for irradiating two kinds of excitation light with different wavelength to the medium, a step for detecting the living cells from the fluorescence intensity obtained, and a step for measuring the pH of the medium by calcg. the intensity ratio between the fluorescence generated by two kinds of excitation light.

IC ICM C12Q001-06

ICS C12M001-34; C12Q001-34; G01N021-64; G01N021-80; G01N033-84

C 9-5 (Biochemical Methods)

Section cross-reference(s): 10

ST living cell microorganism medium pH fluorometry

IT Optical filters

(bandpass; method for detecting living cells in medium and measuring pH of medium)

IT Fluorescent substances

(enzyme substrate; method for detecting living cells in medium and measuring pH of medium)  $\frac{1}{2}$ 

IT Light

 $(\mbox{\bf excitation};$  method for detecting living cells in medium and measuring pH of medium)

IT Cell

(living; method for detecting living cells in medium and measuring pH of medium)

IT Aeolosoma

Bacteria (Eubacteria)

Chlorella

Culture media

Cyclidium (protozoan)

Fluorescence microscopy

Fluorometry

Lepadella Microorganism

Philodina

Scenedesmus

Schizothrix

Test kits

Wavelength

На

(method for detecting living cells in medium and measuring pH of

medium)

- L17 ANSWER 20 OF 179 CA COPYRIGHT 2003 ACS
- AN 136:337178 CA
- TI Spectrofluorometric system devised so as to permit the synchronous measurement of both the bacterial bioluminescence and its related **fluorescence emission**
- AU Karatani, Hajime; Furuta, Kenji; Hirayama, Satoshi
- CS Dept. of Polymer Science and Engineering, Kyoto Institute of Technology, Japan
- SO Proceedings of SPIE-The International Society for Optical Engineering (2001), 4252 (Advances in Fluorescence Sensing Technology V), 88-96 CODEN: PSISDG; ISSN: 0277-786X
- PB SPIE-The International Society for Optical Engineering
- DT Journal
- LA English
- A novel spectrofluorometer equipped with two photo-detectors has been AB developed based on a conventional spectrofluorometer with a view to measuring both bioluminescence and its related fluorescence originating from bacterial luciferase reaction. The configuration of the two monochromator-photomultiplier systems is to be opposite to each other. First the capability of the created system was evaluated using the peroxidase-catalyzed luminol chemiluminescence. The light emission signals as a function of time evidently showed that intensities of chemiluminescence generated by the reaction and of fluorescence elicited from the photoexcited reaction product vary inversely. This reflects the feature of the luminol reaction. Any interference from either excitation light or fluorescence emission is absent in the detection of chem. initiated light. Subsequently, the evaluation of the spectrofluorometer was made on the luciferase reactions under the various conditions. The observation of signals for bioluminescence and fluorescence from the luciferase reactions under the various conditions. The observation of signals for bioluminescence and **fluorescence** from the luciferase reaction was established to be useful in studying the time-dependent behaviors of the fluorescent substrates and/or products as well as a primary emitter formed in the luciferase reaction. From these evaluations, the developed spectrofluoromenter has proved to be profitable to study bacterial bioluminescence.
- CC 9-5 (Biochemical Methods)

Section cross-reference(s): 10

- ST spectrofluorometry measurement bacteria bioluminescence fluorescence
- IT Bacteria (Eubacteria)

## Fluorescence

Fluorometry

Luminescence, bioluminescence

(spectrofluorometric system for synchronous measurement of both bacterial bioluminescence and related **fluorescence emission**)

- IT 9014-00-0, Luciferase 39346-42-4, FMN reductase
  - RL: ANT (Analyte); ANST (Analytical study)

(spectrofluorometric system for synchronous measurement of both bacterial bioluminescence and related **fluorescence** 

emission)

RE.CNT 12 THERE ARE 12 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

```
ANSWER 33 OF 179 CA COPYRIGHT 2003 ACS
L17
AN
     135:192324 CA
     Ultraviolet fluorescence imaging applications
ΤI
     Hill, Ralph H., Jr.; Angell, Peter
AU
     Instrumentation and Space Research Division, Southwest Research Institute,
CS
     San Antonio, TX, USA
     AT-PROCESS (2000), 5(3,4), 108-114
SO
     CODEN: APJCFR; ISSN: 1077-419X
PB
     InfoScience Services
DT
     Journal
LA
    English
    The UV fluorescence of arom. amino acids in microbial
AB
     biofilms can be used to det. the biomass formed in corrosion pits on metal
     surfaces. This information is important in establishing the relationship
     between bacteria and corrosion; i.e., which comes first, the
     corrosion pits or the biomass One specific amino acid that has been used
     in past studies to indicate biomass is tryptophane. Tryptophane
     fluoresces in the near UV region; this fluorescence can be used
     to quantify the amt. of tryptophane and biomass, present on the metal
     surfaces. Under Southwest Research Institute (SwRI) internal research
     funding, a radiometrically calibrated UV-imaging system has been
     previously developed. This system was developed to image tactical missile
     plumes in the solar-blind UV region of the spectrum near 260 nm. The
     basic building block is a microchannel plate-intensified charge-coupled
     device (MCP-CCD) camera. For the project presented in this paper, a 325
     nm helium-cadmium laser was used as an excitation source and the
     camera filtering converted to the near-UV wavelength region. Research was
     conducted to measure the fluorescence from the arom. amino acid
     tryptophane under various dilns. in water. Fluorescent images were also
     recorded from the Oceanospirillum bacteria on a copper coupon;
     this bacteria was originally isolated from copper on Navy
     platforms by Naval Research Lab. personnel. Other applications will also
     be mentioned.
CC
     9-5 (Biochemical Methods)
ST
     UV fluorescence imaging
TT
     Lasers
        (Helium-cadmium; UV fluorescence imaging applications)
     Bacteria (Eubacteria)
TT
     Biofilms (microbial reactors)
     Riomass
     CCD cameras
     Corrosion
     Dilution
       Fluorescence
     Interface
     Oceanospirillum
        (UV fluorescence imaging applications)
     Metals, biological studies
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (UV fluorescence imaging applications)
TΤ
     Imaging
        (UV fluorescence; UV fluorescence imaging
        applications)
     Amino acids, analysis
ΙT
     RL: ANT (Analyte); BSU (Biological study, unclassified); ANST
     (Analytical study); BIOL (Biological study)
        (arom.; UV fluorescence imaging applications)
ΙT
     Wavelength
        (near-UV; UV fluorescence imaging applications)
     73-22-3, Tryptophane, analysis
IT
     RL: ANT (Analyte); BSU (Biological study, unclassified); ANST
     (Analytical study); BIOL (Biological study)
        (UV fluorescence imaging applications)
```

7440-50-8, Copper, biological studies 7732-18-5, Water, biological IT studies RL: BSU (Biological study, unclassified); BIOL (Biological study) (UV fluorescence imaging applications) THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT ANSWER 36 OF 179 CA COPYRIGHT 2003 ACS 135:18679 CA AN Association of bright greenish yellow fluorescence with ΤI aflatoxin production in cereals Ijaz, Nusrat; Salah-ud-Din; Yasin, M. AU Biotechnology and Food Research Centre, Pakistan Council of Scientific and CS Industrial Research Laboratories, Lahore, Pak. SO Pakistan Journal of Science (2000), 52(1-2), 47-52 CODEN: PAJSAS; ISSN: 0030-9877 Pakistan Association for the Advancement of Science PΒ DT Journal English LA AB Bright greenish yellow fluorescence (BGYF) was studied in cereals, after inoculating those with spores of toxic strain of Aspergillus flavus. Pos. relationship between mold growth, kojic acid prodn., BGYF emission and aflatoxin contents was established. BGYF units increased with an increase in aflatoxin prodn. Profused mold growth resulted in improved synthesis of Kojic acid, the chem. which is necessary to produce BGYF. CC17-1 (Food and Feed Chemistry) ST cereal aflatoxin fluorescence fluorimeter TΤ Aspergillus flavus Cereal (grain) Fluorescence Fluorometers Rice (Oryza sativa) Sorghum Wheat (assocn. of bright greenish yellow fluorescence with aflatoxin prodn. in cereals) Aflatoxins RL: ANT (Analyte); BOC (Biological occurrence); BSU (Biological study, unclassified); ANST (Analytical study); BIOL (Biological study); OCCU (Occurrence) (assocn. of bright greenish yellow fluorescence with aflatoxin prodn. in cereals) 9003-99-0, Peroxidase 501-30-4, Kojic acid TT RL: BOC (Biological occurrence); BSU (Biological study, unclassified); BIOL (Biological study); OCCU (Occurrence) (assocn. of bright greenish yellow fluorescence with aflatoxin prodn. in cereals) THERE ARE 17 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT L17 ANSWER 42 OF 179 CA COPYRIGHT 2003 ACS ΑN 134:233815 CA ΤI Physical perturbation for fluorescent characterization of microorganism particles ΑU Bronk, Burt V.; Shoaibi, Azadeh; Nudelman, Raphael; Akinyemi, Agnes N. AFRL/ECBC at U.S. Army ECBC, A.P.G., MD, 21010-5424, USA CS SO Proceedings of SPIE-The International Society for Optical Engineering (2000), 4036 (Chemical and Biological Sensing), 169-180 CODEN: PSISDG; ISSN: 0277-786X PΒ SPIE-The International Society for Optical Engineering DT Journal LA English The motivation for using response to phys. perturbation to classify

microparticles came from our previous expts. With Dipicolinic Acid (DPA).

DPA as a calcium complex is a major component of bacterial spores, constituting more than 5% of their dry wt. It is not commonly found in other natural products and therefore its presence is indicative of the presence of bacterial spores. Previous schemes utilizing the presence of DPA to detect these spores have relied on fluorescence which occurs when lanthanide metals (e.g., terbium) are added to a soln, where the presence of DPA is to be detd. We have recently demonstrated that changes in the fluorescence of DPA can be stimulated without the addn. of such reagents. Thus after exposure to UV light, a substantial increase of fluorescence emitted by DPA solns, with a peak at 410 nm occurs for excitation light with wavelength less than approx. 305 nm.

P.5. (Biochemical Methods)

CC 9-5 (Biochemical Methods)
Section cross-reference(s): 10

ST fluorometry microorganism optical classification dipicolinate

IT Bacillus megaterium
Bacillus subtilis
Escherichia coli
Fluorometry
Microorganism

(phys. perturbation for fluorescent characterization of microorganism particles)

IT 499-83-2, Dipicolinic Acid

RL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses)

(phys. perturbation for fluorescent characterization of microorganism particles)

RE.CNT 11 THERE ARE 11 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L17 ANSWER 46 OF 179 CA COPYRIGHT 2003 ACS

AN 134:128009 CA

TI Sensitivity of detection of **bacteria** with fluorescent and luminescent phenotypes using different instruments

AU Brovko, Lioubov Yu.; Griffiths, Mansel W.

CS Food Sci. Dep., Univ. of Guelph, Guelph, ON, Can.

- Proceedings of SPIE-The International Society for Optical Engineering (2000), 3921(Optical Diagnostics of Living Cells III), 147-156 CODEN: PSISDG; ISSN: 0277-786X
- PB SPIE-The International Society for Optical Engineering

DT Journal

LA English

The problem of bacterial enumeration in different samples is of great AB importance in many fields of research. Construction of recombinant fluorescent and luminescent bacteria that can be easily detected by nondestructive instrumental methods provides us with an opportunity to monitor bacteria in a wide variety of clin., environmental and food samples in real time. Three different labels were employed: Green Fluorescent Protein (GFP), Bacterial luciferase (BL) and Firefly Luciferase (FFL). Both plasmid and chromosomal transformants of different strains of E. coli, P. putida and S. enteritidis were used. For the detection of the in vivo GFP the Shimadzu RF 540 spectrofluorimeter, Labsystems FL- 500 plate fluorometer and Night Owl LB 98 CCD-camera from EG and G Berthold supplied with excitation light source and proper spectral filters both in macroscopic and microscopic mode were used. For the detection of in vivo luminescence of BL and FFL, tube luminometer BG-P from GEM Biomedical Inc., luminometric plate reader from BioOrbit, BIQ Bioview CCD camera from Cambridge Imaging Ltd and Night Owl LB 98 CCD camera both in macroscopic and microscopic mode were used. The expression levels of the labels, their stability, stability of the signal and detection limits of tagged bacteria were investigated. The detection limits for GFP tagged bacteria were 5 X 104 - 6 X 106, for BL tagged bacteria 5 X 102 - 2 X 105, and for FFL tagged

bacteria - 4 X 103 - 106 CFU/mL, depending on the instrument used. Single bacteria could be detected with the help of the Night Owl in the microscopic mode. 9-5 (Biochemical Methods) Section cross-reference(s): 10 ST bacteria enumeration fluorescence luminescence luciferase GFP app Luminescence spectroscopy ΙT (bioluminescence; sensitivity of detection of bacteria with fluorescent and luminescent phenotypes using different instruments) Proteins, specific or class TT RL: ANT (Analyte); ARU (Analytical role, unclassified); BPR (Biological process); BSU (Biological study, unclassified); BUU (Biological use, unclassified); ANST (Analytical study); BIOL (Biological study); PROC (Process); USES (Uses) (green fluorescent; sensitivity of detection of bacteria with fluorescent and luminescent phenotypes using different instruments) ΙT Apparatus (luminometer; sensitivity of detection of bacteria with fluorescent and luminescent phenotypes using different instruments) ΙT Bacteria (Eubacteria) CCD cameras Escherichia coli Fluorescence Fluorometers Fluorometry Luminescence Luminescence, bioluminescence Luminescence spectroscopy Pseudomonas putida Salmonella enteritidis (sensitivity of detection of bacteria with fluorescent and luminescent phenotypes using different instruments) 9014-00-0, Bacterial luciferase 61970-00-1, Firefly Luciferase RL: ANT (Analyte); ARU (Analytical role, unclassified); BPR (Biological process); BSU (Biological study, unclassified); BUU (Biological use, unclassified); ANST (Analytical study); BIOL (Biological study); PROC (Process); USES (Uses) (sensitivity of detection of bacteria with fluorescent and luminescent phenotypes using different instruments) THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD RE.CNT 9 ALL CITATIONS AVAILABLE IN THE RE FORMAT L17 ANSWER 47 OF 179 CA COPYRIGHT 2003 ACS AN 134:112423 CA Multispectral bacterial identification ΤI Tanner, Michael A.; Coleman, William J.; Everett, Christine L.; Robles, ΑU Steven J.; Dilworth, Michael R.; Yang, Mary M.; Youvan, Douglas C. CS Kairos Scientific, Inc., Santa Clara, CA, USA Proceedings of SPIE-The International Society for Optical Engineering SO (2000), 3913(In-Vitro Diagnostic Instrumentation), 45-53 CODEN: PSISDG; ISSN: 0277-786X SPIE-The International Society for Optical Engineering PΒ DT Journal LA English A multi spectral optical technique was developed to simultaneously AΒ classify individual bacterial cells within mixed populations. Multi spectral Bacterial Identification (mBID) combines innovations in microscopy with a software anal. program to measure and characterize the fluorescence signals from multiplexed 16S rRNA probes hybridized to populations of different bacteria. Software was developed to identify individual bacteria at the level of species within these mixed populations. TO test the feasibility of mBID, we examd. the fluorescence emissions from a mixt. of probes specific

for individual species of known bacteria from the American Type Culture Collection. Currently, up to seven species can be detected simultaneously by fluorescence microscopy. An eighth signal was reserved for a universal probe to control for fluorescence intensity. MBID can also be used to identify uncultured microorganisms. We plan to couple this new multi spectral technol. to existing identification technologies that utilize 16S rRNA sequence alignment. Using this integrated identification protocol, bacteria that may be assocd. with chronic conditions will be identified first by analyzing their 16S rDNA sequences and then by visualizing them with fluorescent probes hybridized to their 16S rRNA in situ. 9-4 (Biochemical Methods) Section cross-reference(s): 10, 14 multispectral bacterial identification fluorescence microscopy rRNA specific probe vaginosis rRNA RL: ARG (Analytical reagent use); ANST (Analytical study); USES (16 S; multispectral bacterial identification) Vagina (disease; multispectral bacterial identification) Arthrobacter oxidans Bacillus subtilis Bacteria (Eubacteria) Corynebacterium flavescens Escherichia coli Flexibacter maritimus Fluorescence microscopy Lactobacillus delbrueckii lactis Leptothrix discophora Nucleic acid hybridization (multispectral bacterial identification) 312978-93-1D, conjugate with Bodipy 564/570 PL: ARG (Analytical reagent use); ANST (Analytical study); USES (Arthrobacter globiformis 16S rRNA specific probe; multispectral bacterial identification) 312978-92-0D, conjugate with Bodipy R6G PL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses) (Bacillus subtilis 16S rRNA specific probe; multispectral bacterial identification) 312978-95-3D, conjugate with Cy5 RL: ARG (Analytical reagent use); ANST (Analytical study); USES (Corynebacterium flavescens 16S rRNA specific probe; multispectral bacterial identification) 312978-91-9D, conjugate with Pacific Blue PL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses) (Escherichia coli 16S rRNA specific probe; multispectral bacterial identification) 312978-96-4D, conjugate with Cy5.5 RL: ARG (Analytical reagent use); ANST (Analytical study); USES (Flexibacter maritimus 16S rRNA specific probe; multispectral bacterial identification) 312978-90-8D, conjugate with Alexa 350 PL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses) (Lactobacillus lactis 16S rRNA specific probe; multispectral bacterial identification) 312978-94-2D, conjugate with Bodipy 581/591 FL: ARG (Analytical reagent use); ANST (Analytical study); USES

CC

ST

ΙT

ΙT

IT

IT

IT

TТ

IT

IT

IT

ΙT

(Uses) (Leptothrix discophora 16S rRNA specific probe; multispectral bacterial identification) 320797-24-8D, conjugate with Bodipy 493/503 RL: ARG (Analytical reagent use); ANST (Analytical study); USES (bacterial 16S rRNA specific probe; multispectral bacterial identification) 989-38-8D, conjugate with oligonucleotide probe 121207-31-6D, Bodipy 493/503, conjugate with oligonucleotide probe 144377-05-9D, conjugate with oligonucleotide probe 150152-69-5D, Bodipy 581/591, conjugate with oligonucleotide probe 150173-89-0D, Bodipy 564/570, conjugate with 172777-84-3D, Cy5.5, conjugate with oligonucleotide probe 215868-31-8D, Pacific Blue, conjugate with oligonucleotide probe oligonucleotide probe 244636-14-4D, Alexa 350, conjugate with oligonucleotide probe RL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses) (multispectral bacterial identification) RE.CNT 28 THERE ARE 28 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT ANSWER 50 OF 179 CA COPYRIGHT 2003 ACS L17 133:198214 CA ANUse of fluorescence for characterizing source and speciation of TT aquatic humic substances McKnight, Diane M.; Klapper, Lisa; Hood, Eran W.; Boyer, Elizabeth W. ΑU Institute of Arctic and Alpine Research, University of Colorado, Boulder, CS CO, 80309, USA Preprints of Extended Abstracts presented at the ACS National Meeting, SO American Chemical Society, Division of Environmental Chemistry (2000), 40(2), 659-660 CODEN: PEACF2; ISSN: 1524-6434 American Chemical Society, Division of Environmental Chemistry PB DT Journal LA English Aq. humic substances are defined as a heterogeneous class of moderate mol. AB wt., yellow-colored bio-mols. which are present in all natural water. Spectroscopic characterization of humic substances provides only limited structural information due to the heterogeneity within fulvic acids, although spectroscopic characterization is useful in quantifying differences among fulvic acids. Fluorescence, which can be detected at fulvic acid concns. present in most natural water, potentially contains information about the source and speciation of humic substances. Humic substances derived from degrdn. of microbial matter have distinctive fluorescence characteristics vs. those derived from plant and soil matter. Based on differences in representative excitation emission matrixes, a simple index was developed which could be used in a field study involving a large no. of samples. This index is the ratio of emission at 450 nm to the emission at 500 nm for an excitation of 370 nm. This index has a value of .apprx.1.8 for microbially-derived fulvic acids and .apprx.1.25 for terrestrially-derived fulvic acids. Since other chem. characteristics of fulvic acid vary between these 2 sources of org. matter, these results may indicate a significant seasonal change in fulvic acid reactivity. Thus, fluorescence measurements may provide a tool to est. humic substance reactivity in different environments and provide ancillary data to understand C cycling in aq. ecosystems. CC 61-3 (Water) Section cross-reference(s): 80 aquatic humic substance speciation source characterization; ST fluorescence characterization aquatic humic substance ΙT Fulvic acids Humus

RL: ANT (Analyte); OCU (Occurrence, unclassified); ANST (Analytical study); OCCU (Occurrence)

(aq.; using **fluorescence** to characterize sources and speciation of aquatic humic substances)

IT Fluorescence

(using **fluorescence** to characterize sources and speciation of aquatic humic substances)

IT 7732-18-5, Water, analysis

RL: AMX (Analytical matrix); ANST (Analytical study) (natural water; using fluorescence to characterize sources and speciation of aquatic humic substances)

RE.CNT 9 THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L17 ANSWER 56 OF 179 CA COPYRIGHT 2003 ACS

AN 132:185144 CA

- TI A novel method for detection of viable Giardia cysts in water samples
- AU Jarmey-Swan, C.; Gibbs, R. A.; Ho, G. E.; Bailey, I. W.; Howgrave-Graham, A. R.
- CS Analytical Services, Umgeni Water, Pietermaritzburg, 3200, S. Afr.
- SO Water Research (2000), 34(6), 1948-1951 CODEN: WATRAG; ISSN: 0043-1354
- PB Elsevier Science Ltd.
- DT Journal
- LA English
- Assessing Giardia viability is a major requirement for public health AB purveyors and the water industry. Several indicators of viability (e.g., stains, excystation, animal infectivity) have been used to enumerate cysts with varying degrees of success. A combined detection-viability method for use in water samples would be useful to detect and det. cyst viability in source and drinking water and disinfection efficacy at treatment plants. Distd. water samples were seeded with purified Giardia cysts and incubated with fluorescein diacetate (FDA), initially to stain viable cysts, followed by tetra-Methyl red labeled anti-Giardia monoclonal antibodies (TMR) for confirmation of identity. As a result of FDA staining, green fluorescence of intact viable cysts was obsd. microscopically using a 450-490 nm exciter filter; non-viable cysts were not stained. Giardia cysts reacted pos. with TMR and glowed red using a triple band microscope filter with excitations of 400/450/570nm. At this wavelength, a combination of FDA and TMR stained viable cysts green internally with a red wall while non-viable cysts only stained red. This simple, reliable, quick method allowed differentiation of Giardia cysts in water samples while simultaneously detg. their viability.
- CC 61-3 (Water)

Section cross-reference(s): 10, 80

- ST Giardia cyst viability detn water; staining monoclonal antibody identification viability confirmation Giardia cyst; fluorescein diacetate staining Giardia cyst water; tetramethylrhodamine labeled monoclonal antibody Giardia cyst identification
- IT Cyst, microbial

(Giardia; staining/monoclonal antibody method to simultaneously detect, identify, and confirm viability of Giardia cysts in water)

IT Giardia

(cysts; staining/monoclonal antibody method to simultaneously detect, identify, and confirm viability of Giardia cysts in water)

IT Antibodies

RL: ARG (Analytical reagent use); NUU (Other use, unclassified); ANST (Analytical study); USES (Uses)

(monoclonal, tetramethylrhodamine-labeled; staining/monoclonal antibody method to simultaneously detect, identify, and confirm viability of Giardia cysts in water)

IT 596-09-8, Fluorescein diacetate

RL: ARG (Analytical reagent use); NUU (Other use, unclassified); ANST (Analytical study); USES (Uses)

(cyst staining with; staining/monoclonal antibody method to simultaneously detect, identify, and confirm viability of Giardia cysts in water)

IT 70281-37-7, Tetramethylrhodamine

RL: ARG (Analytical reagent use); NUU (Other use, unclassified); ANST

(Analytical study); USES (Uses)

(monoclonal antibodies labeled with; staining/monoclonal antibody method to simultaneously detect, identify, and confirm viability of Giardia cysts in water)

IT 7732-18-5, Water, analysis

RL: AMX (Analytical matrix); ANST (Analytical study) (source water and drinking water; staining/monoclonal antibody method to simultaneously detect, identify, and confirm viability of Giardia cysts in water)

RE.CNT 9 THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L17 ANSWER 60 OF 179 CA COPYRIGHT 2003 ACS
- AN 131:141627 CA
- TI Fluorescence of dipicolinic acid as a possible component of the observed UV emission spectra of bacterial spores
- AU Nudelman, Raphael; Feay, Nicole; Hirsch, Mathew; Efrima, Shlomo; Bronk, Burt
- CS Mantech Environmental Technology Inc., USA
- SO Proceedings of SPIE-The International Society for Optical Engineering (1999), 3533(Air Monitoring and Detection of Chemical and Biological Agents), 190-195
  CODEN: PSISDG; ISSN: 0277-786X
- PB SPIE-The International Society for Optical Engineering
- DT Journal
- LA English
- Dipicolinic acid (DPA) and the Ca2+ complex of DPA (CaDPA) are well-known and are major chem. components of bacterial spores. DPA's native fluorescence is very weak and is thought to be completely masked by the fluorescence of tryptophan when this compd. is present. Thus fluorescence related to DPA in spores is assumed by many authors to be completely absent. We show that the fluorescence of CaDPA is substantial for excitation between about 290 nm and 310 nm with emission peaking near 400 nm. This emission is at the long wavelength tail for emission from tryptophan. We examine whether the emission of CaDPA could contribute to the total emission spectrum when bacterial spores are present in an aerosol, for excitation wavelengths in the neighborhood of 310 nm. In this report we present measurements of fluorescence excitation and emission for CaDPA and compare them with that of DPA and tryptophan.
- CC 9-5 (Biochemical Methods)

Section cross-reference(s): 4, 10

- ST bacterium spore dipicolinate fluorometry
- IT Bacteria (Eubacteria)
  Environmental analysis
  Fluorometry

Spore

(fluorescence of dipicolinic acid as a possible component of obsd. UV emission spectra of bacterial spores)

TT 73-22-3, L-Tryptophan, analysis 499-83-2, Dipicolinic acid 499-83-2D, Dipicolinic acid, calcium complex 7440-70-2D, Calcium, complex with dipicolinic acid, analysis
RL: ANT (Analyte); BOC (Biological occurrence); BSU (Biological study, unclassified); ANST (Analytical study); BIOL (Biological study); OCCU (Occurrence)

(fluorescence of dipicolinic acid as a possible component of obsd. UV emission spectra of bacterial spores)

## RE.CNT 15 THERE ARE 15 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L17 ANSWER 64 OF 179 CA COPYRIGHT 2003 ACS

AN 130:278941 CA

TI Fluorescent biological particle detection system

IN Ho, Jim Yew-wah

PA Her Majesty the Queen In Right of Canada, as Represented by the Minister of, Can.

SO U.S., 18 pp., Cont.-in-part of U.S. 5,701,012.

CODEN: USXXAM

DT Patent

LA English

FAN.CNT 2

|      | PATENT NO.     | KIND | DATE     | APPLICATION NO. | DATE     |
|------|----------------|------|----------|-----------------|----------|
|      |                |      |          |                 |          |
| ΡI   | US 5895922     | A    | 19990420 | US 1997-863023  | 19970523 |
|      | US 5701012     | Α    | 19971223 | US 1996-616475  | 19960319 |
| PRAI | US 1996-616475 |      | 19960319 |                 |          |

AB A process and app. are provided for detection of viable and potentially hazardous biol. particles which may be dispersed in a particulate-contg. airstream. The process comprises directing each of the contained particles along a linear path through air, in a sequential manner, and sampling them for detn. of their size, whether they are biol. and viable, and whether they are present in concns. greater than background levels. The particle size identifies the particles as respirable or not and the particles are characterized as biol. and viable by subjecting each particle in turn, to 340 nm, UV laser light and looking for the

emission of fluorescence which is typically emitted from

bacteria or bacterial spore. Fluorescence detected in the 400-540 nm range signals the presence of NADH, which is indicative of biol. activity or viability. A more compact, and power-saving app. results with the preferential use of a solid state, UV laser, which is actuated only when the particle is passing the laser and only if it is deemed to be a biol. viable candidate.

IC ICM G01N021-64

NCL 250491200

CC 9-5 (Biochemical Methods)

ST fluorescence biol particle detection system

IT Particles

(Biol.; fluorescent biol. particle detection system)

IT Air analysis

Apparatus

Bacteria (Eubacteria)

Biochemical molecules

Fluorometry

Particle size

Sampling

Spore

UV lasers

(fluorescent biol. particle detection system)

IT 58-68-4, NADH

RL: ANT (Analyte); BSU (Biological study, unclassified); ANST

(Analytical study); BIOL (Biological study)

(fluorescent biol. particle detection system)

RE.CNT 10 THERE ARE 10 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L17 ANSWER 67 OF 179 CA COPYRIGHT 2003 ACS
- AN 130:179461 CA
- TI Two-photon excitation in fluorescence lifetime imaging
- AU Gerritsen, Hans C.; Vroom, Jurrien; Sytsma, Joost
- CS Debye Institute, Utrecht University, Utrecht, 3508 TA, Neth.
- SO Fluorescence Microscopy and Fluorescent Probes, [Based on the Proceedings

```
of the Conference on Fluorescence Microscopy and Fluorescent Probes], 2nd,
    Prague, Apr. 9-12, 1997 (1998), Meeting Date 1997, 55-62. Editor(s):
    Slavik, Jan. Publisher: Plenum, New York, N. Y.
    CODEN: 67BTAH
DT
    Conference
    English
LA
    The authors employ time-gated lifetime imaging implemented in a two-photon
AB
    excitation scanning microscope. The examples given demonstrate
    that fluorescence lifetime contrast is particularly suitable for
    in-depth imaging expts. Lifetime imaging enables the discrimination of
    multiple probes based on their differences in lifetime. The technol. is
    applied to the imaging of biofilms and human skin using Acridine-Orange.
    9-4 (Biochemical Methods)
CC
    Section cross-reference(s): 6, 13
    microscopy two photon excitation fluorescence lifetime
ST
    imaging
    Staining, biological
ΙT
        (fluorescent; two-photon excitation in fluorescence
       lifetime imaging)
ΙT
    Tooth
        (plaque; two-photon excitation in fluorescence
       lifetime imaging)
ΙΤ
    Imaging
        (time-gated lifetime; two-photon excitation in
       fluorescence lifetime imaging)
TT
    Biofilm bacteria
    Fluorescent indicators
    Fluorescent substances
        (two-photon excitation in fluorescence lifetime
       imaging)
TΤ
    Microscopy
        (two-photon excitation scanning; two-photon
       excitation in fluorescence lifetime imaging)
    Laser induced fluorescence
IT
    Photoexcitation
        (two-photon; two-photon excitation in fluorescence
       lifetime imaging)
ΙT
    65-61-2, Acridine Orange
    RL: ARU (Analytical role, unclassified); BSU (Biological study,
    unclassified); PRP (Properties); ANST (Analytical study); BIOL
     (Biological study)
        (two-photon excitation in fluorescence lifetime
       imaging)
             THERE ARE 18 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT 18
             ALL CITATIONS AVAILABLE IN THE RE FORMAT
L17 ANSWER 79 OF 179 CA COPYRIGHT 2003 ACS
AN
    128:45574 CA
ΤI
    Fluorescent biological particle detection system
IN
    Ho, Jim Yew-wah
    Her Majesty the Queen In Right of Canada, as Represented by the Minister
PA
    of, Can.
SO
    U.S., 17 pp.
    CODEN: USXXAM
DT
    Patent
    English
LA
FAN.CNT 2
                   KIND DATE
                                        APPLICATION NO. DATE
    PATENT NO.
    ______
    US 5701012 A 19971223
                                         US 1996-616475
                                                         19960319
PΤ
    US 5895922
                    A 19990420
                                         US 1997-863023
                                                         19970523
PRAI US 1996-616475
                           19960319
AB A process and app. are provided for detection of viable and potentially
```

hazardous biol. particles which may be dispersed in a particulate-contg. airstream. The process comprises directing each of the contained particles along a linear path through air, in a sequential manner, and sampling them for detn. of their size, whether they are biol. and viable, and whether they are present in concns. greater than background levels. The particle size identifies the particles as respirable or not and the particles are characterized as biol. and viable by subjecting each particle in turn, to 340 nm, UV laser light and looking for the emission of fluorescence which is typically emitted from bacteria or bacterial spore. Fluorescence detected in the 400-540 nm range signals the presence of NAD hydrogen, which is indicative of biol. activity or viability. IC ICM G01N021-64 NCL 250461200 9-5 (Biochemical Methods) CCSection cross-reference(s): 10 ST fluorescence biol particle detection system ΙT Spore (Bacterial; fluorescent biol. particle detection system) ΙT Particles (Biol.; fluorescent biol. particle detection system) IT Air analysis Apparatus Bacteria (Eubacteria) Biochemical molecules Fluorometers Fluorometry UV lasers (fluorescent biol. particle detection system) 58-68-4, NADh IT RL: ANT (Analyte); ANST (Analytical study) (fluorescent biol. particle detection system) L17 ANSWER 88 OF 179 CA COPYRIGHT 2003 ACS AN127:80177 CA Noninvasive monitoring of the physiological state of microbial ΤI cultures ΑU Lestan, D.; Perdih, A. Centre Soil Environmental Sci., Dep. Agronomy, Biotechnical Faculty, Univ. CS Ljubljana, Ljubljana, Slovenia Acta Chimica Slovenica (1997), 44(1), 1-15 SO CODEN: ACSLE7; ISSN: 1318-0207 Slovenian Chemical Society PB Journal; General Review DTLA English A review with 41 refs. In attempts to improve the performance of ΔR bioprocess modeling and control it is becoming clear that alternative methods for accessing information from biol. systems, better suited to the nature of living systems, have to be developed. Three considerably different approaches have been proposed to access this information and are reviewed here. The optical approach relies on enzymes or metabolites which change their optical absorption or fluorescence emission as a function of specific or induced cellular alterations. 31P NMR can be used for the detn. of energy-rich P compds. Online monitoring of the physiol. state of the living matter in bioreactors uses knowledge-based recognition systems to assess variables that indicate the physiol. state. 16-0 (Fermentation and Bioindustrial Chemistry) CCSection cross-reference(s): 9, 10 review microorganism culture noninvasive monitoring; optical spectroscopy ST microbial culture monitoring review; phosphorus 31 NMR culture monitoring review; recognition system microbial culture monitoring review ΙT Computer application

```
rexpert systems; noninvasive monitoring of physiol. state of
       microbial cultures)
IT
     Bioreactors
    Biotechnology
       Fluorescence
     Microorganism
     Optical absorption
     Spectroscopy
     UV and visible spectroscopy
        (noninvasive monitoring of physiol. state of microbial
        cultures)
ΙT
     Energy-rich phosphates
     RL: ANT (Analyte); ANST (Analytical study)
        (noninvasive monitoring of physiol. state of microbial
        cultures)
    NMR spectroscopy
TΤ
        (phosphorus-31; noninvasive monitoring of physiol. state of
        microbial cultures)
L17
    ANSWER 98 OF 179 CA COPYRIGHT 2003 ACS
AN
     124:64699 CA
     Aerosol-fluorescence spectrum analyzer: real-time measurement of
ΤI
     emission spectra of airborne biological particles
     Hill, Steven C.; Pinnick, Ronald G.; Nachman, Paul; Chen, Gang; Chang,
ΑU
     Richard K.; Mayo, Michael W.; Fernandez, Gilbert L.
     Army Research Laboratory, White Sands Missile Range, NM, 88002-5501, USA
CS
     Applied Optics (1995), 34(30), 7149-55
SO
     CODEN: APOPAI; ISSN: 0003-6935
PΒ
    Optical Society of America
     Journal
DT
LA
    English
     We have assembled an aerosol-fluorescence spectrum analyzer
AB
     (AFS), which can measure the fluorescence spectra and elastic
     scattering of airborne particles as they flow through a laser beam.
     aerosols traverse a scattering cell where they are illuminated with
     intense (50 kW/cm2) light inside the cavity of an argon-ion laser
     operating at 488 nm. This AFS can obtain fluorescence spectra
     of individual dye-doped polystyrene microspheres as small as 0.5 .mu.m in
    diam. The spectra obtained from microspheres doped with pink and
     green-yellow dyes are clearly different. We have also detected the
     fluorescence spectra of airborne particles (although not single
     particles) made from various biol. materials, e.g., Bacillus subtilis
     spores, Bacillus anthracis spores, riboflavin, and tree
     leaves. The AFS may be useful in detecting and characterizing airborne
    bacteria and other airborne particles of biol. origin.
CC
    59-1 (Air Pollution and Industrial Hygiene)
    airborne biol aerosol particle detn; bioaerosol detn air laser
ST
    fluorescence
    Air analysis
ΙT
     Particles
        (real-time measurement of emission spectra of airborne biol.
       particles)
IT
     Bacillus anthracis
     Bacillus subtilis
        (spores; real-time measurement of emission spectra
       of airborne biol. particles)
TT
    Aerosols
        (airborne, biol., real-time measurement of emission spectra
       of airborne biol. particles)
    Air pollution
ΙT
        (particulate, real-time measurement of emission spectra of
       airborne biol. particles)
ΙT
     83-88-5, Riboflavin, analysis
    RL: ANT (Analyte); ANST (Analytical study)
```

(real-time measurement of **emission** spectra of airborne biol. particles)

```
L17 ANSWER 100 OF 179 CA COPYRIGHT 2003 ACS
```

AN 124:22668 CA

TI Green fluorescent protein as a new expression marker in mycobacteria

AU Kremer, Laurent; Baulard, Alain; Estaquier, Jerome; Poulain-Godefroy, Odile; Locht, Camille

CS Laboratoire de Microbiologie Genetique et Moleculaire, INSERM, Lille Cedex, F-59019, Fr.

SO Molecular Microbiology (1995), 17(5), 913-22 CODEN: MOMIEE; ISSN: 0950-382X

PB Blackwell

DT Journal

LA English

This study describes the use and the advantages of the green fluorescent AB protein (GFP) as a reporter mol. for mycobacteria. The gfp gene from Aequorea victoria was placed under the control of the HSP60 promoter in the shuttle vector pGFM-11. The gfp expression in the recombinant Mycobacterium smeqmatis and BCG was readily detected on agar plates by the development of an intense green fluorescence upon irradn. with long-wave UV light. In mycobacteria contg. a pGFM-11 deriv. that lacks the hsp60 promoter, no fluorescence was obsd. However, this plasmid was successfully used as a promoter-probe vector to identify RCG promoters. The fluorescence emission of GFP in mycobacteria harboring pGFM-11 and grown in liq. media could be quantified by spectrofluorimetry. This allowed for easy assessment of drug susceptibility. As GFP does not require the addn. of substrates or cofactors, the green fluorescent bacilli could be directly obsd. within infected macrophages using fluorescence and laser confocal microscopy, or in tissue sections of infected mice. Finally, infected cells or free-living recombinant mycobacteria could also be analyzed by flow cytometry. The GFP thus appears to be a convenient reporter for mycobacteria, allowing tracing of recombinant mycobacteria, isolation of promoters with interesting properties, in vivo drug testing, and the development of new diagnostic tools.

CC 3-1 (Biochemical Genetics)

Section cross-reference(s): 9, 10

ST green fluorescent protein expression marker mycobacteria

IT Gene, microbial

RL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses)

(gfp, reporter; green fluorescent protein as an expression marker in mycobacteria)

IT Aequorea victoria

Mycobacterium

Mycobacterium BCG

Mycobacterium smegmatis

(green fluorescent protein as an expression marker in mycobacteria)

Plasmid and Episome

(pGFM-11; green fluorescent protein as an expression marker in mycobacteria)

IT Spectrochemical analysis

(fluorometric, green fluorescent protein as an expression marker in mycobacteria)

IT Proteins, specific or class

RL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses)

(green fluorescent, green fluorescent protein as an expression marker in mycobacteria)

IT Genetic element

RL: BUU (Biological use, unclassified); BIOL (Biological study); USES (Uses)

(promoter, hsp60; green fluorescent protein as an expression marker in

L17 ANSWER 105 OF 179 CA COPYRIGHT 2003 ACS AN 123:51436 CA ΤI Spectroscopic properties of tryptophan and bacteria Tang, G. C.; Yang, Y. L.; Huang, Z. Z.; Hua, W.; Zhou, F.; Cosloy, S.; AU Alfano, R. R. City College, City University of New York, New York, NY, 10031, USA CS Proceedings of SPIE-The International Society for Optical Engineering SO (1995), 2387, 169-72 CODEN: PSISDG; ISSN: 0277-786X DT Journal LA English AB Fluorescence spectra of tryptophan and bacteria were measured at different concns. using a Mediscience CD-Scan unit. The emission spectra of tryptophan were obtained using an excitation wavelength at 280 nm. The excitation spectra were obtained at the emission of 340 nm. The min. detectable concn. of tryptophan was 10-8 M. The emission spectra for bacteria were probed at 340 nm. The min. detectable no. of bacteria in a beam of the excitation light was detd. to be about 30. Assuming that the emission band at 340 nm of bacteria comes from tryptophan, the no. of tryptophan per bacterium was estd. to be 108. This approach to det. the no. is almost consistent with that obtained using a wt. method. CC 9-5 (Biochemical Methods) Section cross-reference(s): 10 fluorescence bacteria tryptophan detection STIT Bacteria Escherichia coli Fluorescence (spectroscopic properties of tryptophan and bacteria) IT 54-12-6, DL-Tryptophan RL: ANT (Analyte); PRP (Properties); ANST (Analytical study) (spectroscopic properties of tryptophan and bacteria) ANSWER 110 OF 179 CA COPYRIGHT 2003 ACS 1.17 AN 121:200084 CA ΤI In-vivo fluorescence detection and imaging of porphyrin-producing bacteria in the human skin and in the oral cavity for diagnosis of acne vulgaris, caries, and squamous cell carcinoma Konig, Karsten; Schneckenburger, Herbert; Hemmer, Joerg; Tromberg, Bruce; ΑU Steiner, Rudolf CS Beckman Laser Institute, Irvine, CA, 92715, USA SO Proceedings of SPIE-The International Society for Optical Engineering (1994), 2135 (Advances in Laser and Light Spectroscopy to Diagnose Cancer and Other Diseases), 129-38 CODEN: PSISDG; ISSN: 0277-786X DT Journal English LA AB Certain bacteria are able to synthesize metal-free fluorescent porphyrins and can therefore be detected by sensitive autofluorescence measurements in the red spectral region. The porphyrin-producing bacterium Propionibacterium acnes, which is involved in the pathogenesis of acne vulgaris, was localized in human skin. Spectrally-resolved fluorescence images of bacteria distribution in the face were obtained by a slow-scan CCD camera combined with a tunable liq. crystal filter. The structured autofluorescence of dental caries and dental plaque in the red is caused by oral bacteria, like Bacteroides or Actinomyces odontolyticus. "Caries images" were created by time-gated imaging in the ns-region after ultrashort laser excitation. Time-gated measurements allow the suppression of backscattered light and non-porphyrin autofluorescence. Biopsies of oral

squamous cell carcinoma exhibited red autofluorescence in necrotic regions

and high concns. of the porphyrin-producing bacterium Pseudomonas aeruginosa. These studies suggest that the temporal and spectral characteristics of bacterial autofluorescence can be used in the diagnosis and treatment of a variety of diseases.

CC 9-5 (Biochemical Methods)

Section cross-reference(s): 8, 10, 14

ST fluorescence imaging porphyrin producing bacterium; acne
vulgaris skin fluorescence imaging; dental caries bacterium
fluorescence imaging; oral squamous cell carcinoma
fluorescence imaging

IT Mouth Skin

(in-vivo **fluorescence** detection and imaging of porphyrin-producing **bacteria** in human skin and in oral cavity for diagnosis of acne vulgaris, caries, and squamous cell carcinoma)

Porphyrins
RL: ANT (Analyte); BOC (Biological occurrence); BSU (Biological study, unclassified); MFM (Metabolic formation); ANST (Analytical study); BIOL (Biological study); FORM (Formation, nonpreparative); OCCU (Occurrence)

(in-vivo **fluorescence** detection and imaging of porphyrin-producing **bacteria** in human skin and in oral cavity for diagnosis of acne vulgaris, caries, and squamous cell carcinoma)

IT Bacteria

(porphyrin-producing; in-vivo fluorescence detection and imaging of porphyrin-producing bacteria in human skin and in oral cavity for diagnosis of acne vulgaris, caries, and squamous cell carcinoma)

IT Tooth

(disease, caries, in-vivo **fluorescence** detection and imaging of porphyrin-producing **bacteria** in human skin and in oral cavity for diagnosis of acne vulgaris, caries, and squamous cell carcinoma)

IT Imaging

(fluorescent, in-vivo **fluorescence** detection and imaging of porphyrin-producing **bacteria** in human skin and in oral cavity for diagnosis of acne vulgaris, caries, and squamous cell carcinoma)

Spectrochemical analysis
(fluorometric, in-vivo fluorescence detection and imaging of
porphyrin-producing bacteria in human skin and in oral cavity
for diagnosis of acne vulgaris, caries, and squamous cell carcinoma)

IT Mouth

IT

(neoplasm, squamous cell carcinoma, in-vivo fluorescence detection and imaging of porphyrin-producing bacteria in human skin and in oral cavity for diagnosis of acne vulgaris, caries, and squamous cell carcinoma)

IT Acne

(vulgaris, in-vivo **fluorescence** detection and imaging of porphyrin-producing **bacteria** in human skin and in oral cavity for diagnosis of acne vulgaris, caries, and squamous cell carcinoma)

- L17 ANSWER 114 OF 179 CA COPYRIGHT 2003 ACS
- AN 120:265052 CA
- TI Online, non-destructive biomass determination of bacterial biofilms by fluorometry
- AU Angell, Peter; Arrage, Andrew A.; Mittelman, Marc W.; White, David C.
- CS Cent. Environ. Biotechno., Knoxville, TN, 37932, USA
- SO Journal of Microbiological Methods (1993), 18(4), 317-27 CODEN: JMIMDQ; ISSN: 0167-7012
- DT Journal
- LA English
- AB The lack of online methodol. for the detn. of **microbial** biomass and activity of attached **bacteria** has severely limited the study of biofilm physiol. This study showed that the fluorescent

emission of arom. amino acids in microbial biofilms can be used to det. the biomass formed on 316 stainless steel coupons. Cells resuspended from the substratum were enumerated by viable and acridine orange counts showing correlation coeffs. of 0.77 and 0.98, resp., when compared to the tryptophan fluorescence. Substrata treated with a fluorescent epoxy coating (F-150) showed no fluorescence that could be attributed to the microorganisms. Bioluminescent emission of an actively growing bioluminescent bacterium, Vibrio harveyi, was correlated with acridine orange counts (r2= 0.95) and fluorescence (r2= 0.93). The results of these studies suggest that fluorescence measurements can be used to monitor microbial biomass assocd. with various substrata. Coupled with bioluminescence measurements, this method provides information on both biomass constituents and metabolic activity, and therefore possibly an indicator of sub-lethal toxicity. 9-5 (Biochemical Methods) Section cross-reference(s): 16, 60 bacteria biofilm biomass detn fluorometry Films (bacterial, biomass of, detn. of, by fluorometry) Bacteria (biofilms, biomass detn. in, by fluorometry) Biomass (detn. of, of bacteria biofilms by fluorometry) 53-57-6, NADPH RL: ANST (Analytical study) (detn. of in bacteria biofilm biomass detn.) 58-68-4, NADH 73-22-3, Tryptophan, analysis RL: ANT (Analyte); ANST (Analytical study) (detn. of, in bacteria biofilm biomass detn.) L17 ANSWER 117 OF 179 CA COPYRIGHT 2003 ACS 119:45156 CA Determination of fluorescent substances in microorganism, and its use for determination of microorganism viability Hirotsuji, Junji; Yoshimura, Yumiko; Sugimoto, Masuo; Nakatsugawa, Naoki; Oota, Naomi Mitsubishi Electric Corp, Japan Jpn. Kokai Tokkyo Koho, 20 pp. CODEN: JKXXAF Patent Japanese FAN.CNT 1 KIND DATE APPLICATION NO. DATE PATENT NO. \_\_\_\_ \_\_\_\_\_\_ A2 19930507 JP 05111394 JP 1991-277705 19911024 B2 19990913 JP 2947305 PRAI JP 1991-277705 19911024 The fluorescent substances such as coenzyme F420 (I) in microorganism is detd. by direct irradn. of the microorganism with an excitation wavelength. Based on the fluorescence data, the content and concn. of the fluorescent substances can be calcd. and the viability of the microorganism detd. The method is easy and fast, and does not need to disrupt the cells. Fluorescent detn. of I in methane bacteria and the viability of the bacteria was shown. The result was comparable to that with the prior art. ICM C120001-02 10-6 (Microbial, Algal, and Fungal Biochemistry) fluorescence detn microorganism viability; substance fluorescence detn microorganism Fluorescent substances (fluorescent detn. of, in microorganism for detn. of viability) Microorganism (fluorescent substances in, detn. of, for detn. of microorganism

CC

ST

ΙT

ΙT

IT

IT

ΙT

AΝ

TI

ΤN

PΑ

SO

DT

LA

AB

ΙC

CCST

IT

TT

```
viability)
    Bacteria
TΤ
       (methanogenic, F420 in, detn. of, for detn. of bacteria
       viability)
    64885-97-8, Coenzyme F420
ΙT
    RL: ANT (Analyte); ANST (Analytical study)
        (fluorescence detn. of, in methane bacteria)
    ANSWER 128 OF 179 CA COPYRIGHT 2003 ACS
AN
    115:178859 CA
    Method and reagents for detecting microorganisms
ΤI
    Monget, Daniel
IN
    Biomerieux S. A., Fr.
PA
    Eur. Pat. Appl., 15 pp.
SO
    CODEN: EPXXDW
    Patent
DT
   French
LA
FAN.CNT 1
                   KIND DATE
                                         APPLICATION NO. DATE
    PATENT NO.
                                         ______
     ______
    EP 424293 A1 19910424
EP 424293 B1 19950412
                                        EP 1990-420453 19901018
        R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE
    FR 2653447 A1 19910426 FR 1989-14087
                     B1 19911227
    FR 2653447
                     E 19950415 AT 1990-420453
                                                         19901018
    AT 121138
                                        ES 1990-420453
                     T3 19950616
                                                          19901018
    ES 2071067
                                         CA 1990-2028059 19901019
                     C 20010213
A 19940809
    CA 2028059
US 5336600
                     A 19940809
                                         US 1992-961625 19921016
PRAI FR 1989-14087 A 19891020
US 1990-600919 B3 19901022
    MARPAT 115:178859
OS
    Microorganisms are detected using an aq. reaction media contg. a C source,
AB
    a N source, and a marker whose luminous emission is modified as
     a consequence of development of the microorganism in the reaction media.
     The marker is, e.g. I (R1, R4, R5 = H, F, Cl, Br, alkyl, alkoxy, CO2H,
     amide, cyano; R2, R3 = H, F, Cl, Br, alkyl, alkoxy, carboxylate, CO2H,
     amide, cyano; or R3R2 form an unsatd. ring; X - OH, amine) or its anionic
     form. Six different antibiotics at 2 different concns. were added to
     tubes contg. Mueller Hinton media, glucose, resorufin, and water (pH 7.3).
     Escherichia coli, Pseudomonas aeruginosa, and Staphylococcus aureus (10
     cells/mL) were added and fluoroescence was measured after 18 h incubation
     at 35.degree.. If the microorganism was sensitive to the antibiotic,
     fluorescence was maximal; resistant microorganisms gave total
     extinction of the fluoroescence.
  ICM C12Q001-04
IC
    ICS C12Q001-18; C07D265-38
    9-5 (Biochemical Methods)
CC
     Section cross-reference(s): 10
     microorganism detection resorufin deriv fluorescence; antibiotic
ST
    microorganism sensitivity fluorescence resorufin
    Candida albicans
TΤ
     Citrobacter freundii
     Escherichia coli
     Pseudomonas aeruginosa
     Staphylococcus aureus
     Streptococcus faecalis
        (detection of, by fluorescence assay, resorufin as marker in)
ΙT
    Microorganism
        (detection of, by luminescence assay, resorufin or deriv. as marker in)
ΙT
     Blood analysis
     Body fluid
     Cerebrospinal fluid
     Cosmetics
```

Food analysis Pharmaceutical analysis Urine analysis umicroorganism detection in, by luminescence assay, resorufin or deriv. as marker for) ΙT Antibiotics Fungicides and Fungistats (microorganism sensitivity to, resorufin or deriv. in luminescence assay of) TT Aeromonas hydrophila Candida tropicalis Klebsiella pneumoniae Proteus vulgaris Torulopsis glabrata Vibrio alginolyticus (response of, to different sugars, in fluorescence assay with resorufin as marker, microorganism identification in relation to) ΙT Spectrochemical analysis (fluorometric, microorganism detection by, resorufin in) ΙT Bacteria (intestinal, detection of, by fluorescence assay, resorufin as marker in) Spectrochemical analysis IT (luminescence, microorganism detection by, resorufin or deriv. as marker in) 635-78-9D, Resorufin, anions 635-78-9, Resorufin IT PL: ANST (Analytical study) (as luminescent marker for microorganism detection) TΤ 101490-85-1 RL: ANST (Analytical study) (as substrate for .beta.-glucuronidase of Escherichia Coli detection) 50-99-7, Glucose, analysis 57-50-1, Saccharose, analysis 63-42-3, Lactose 69-79-4 99-20-7, Trehalose 528-50-7, Cellobiose 585-9 IT Melibiose FL: ANST (Analytical study) (microorganism response to, in microorganism identification by resorufin **fluorescence** assay) 61-33-6, properties 114-07-8, Erythromycin ΙT 60-54-8, Tetracycline 35607-66-0 61477-96-1 PL: ANST (Analytical study) (microorganism sensitivity to, resorufin in fluorescence assay of) IT 9001-45-0 PL: ANST (Analytical study) (resorufin-glucuronide as substrate for, for Escherichia coli detection) L17 ANSWER 141 OF 179 CA COPYRIGHT 2003 ACS ANTΙ Fluorescence photometric determination of the coenzyme F420 to monitor anaerobic effluent purification ΑU Kaiser, G.; Frenzel, S.; Mauch, W. Fachgeb. Zuckertechnol., Tech. Univ. Berlin, Berlin, D-1000/65, Fed. Rep. CS Ger. SO Zuckerindustrie (Berlin, Germany) (1988), 113(10), 868-72 CODEN: ZUCKDI; ISSN: 0344-8657 DT Journal German LA A method is proposed for the detn. of coenzyme F420, which is AB characteristic for CH4-forming bacteria, making it possible to monitor anaerobic effluent purifn. by relatively simple means. equipment required is a slightly modified photometer or nephelometer, a table centrifuge, and a membrane filter. By detg. the fluorescence intensity at 470 nm over a wide spectrum of

excitation wavelengths (340-450 nm), the optimum reaction conditions (e.g. pH, solvent, redox state) for fluorimetric detection have been identified and the specificity of the method for coenzyme F420 established. Since for normal plant control it is not necessary to measure the molar concn. of coenzyme F420, the anal. is reduced to the detn. of the relative fluorescence. The latter reflects the state of effluent fermn. Disturbances in the course of fermn. are quickly manifested by a decrease in fluorescence, allowing prompt correction of the effluent purifn. process. CC 7-1 (Enzymes) Section cross-reference(s): 16 ST coenzyme F420 detn fluorescence methanogen fermn IΤ Fermentation (with methanogens, monitoring of, fluorescence method for) ΙT Bacteria (methanogenic, fermn. with, monitoring of, fluorescence method for) IΤ 64885-97-8, Coenzyme F420 RL: ANT (Analyte); ANST (Analytical study) (detn. of, by fluorescence method, for monitoring fermn. by methanogenic bacteria) ANSWER 149 OF 179 CA COPYRIGHT 2003 ACS L17 AN 105:130360 CA Determination of a concentration of bacteria in a suspension ΤI IN Kosarev, N. V.; Puckkov, E. O. All-Union Scientific-Research Institute of Applied Microbiology, USSR PΑ SÓ U.S.S.R. From: Otkrytiya, Izobret. 1986, (18), 127. CODEN: URXXAF DT Patent LARussian FAN.CNT 1 PATENT NO. KIND DATE APPLICATION NO. DATE ---------\_\_\_\_\_ SU 1231077 A1 19860515 SU 1984-3719309 19840328 PRAI SU 1984-3719309 19840328 The concn. of bacteria in a suspension is detd. by prepg. a bacterial suspension and recording the optical parameters. The accuracy and sensitivity of anal. are increased by adding up to 0.03-0.05% Triton X100, .ltoreq.0.3-3.0 mmole tris-ethylenediaminetetraacetate (sic) at pH 7.2-7.5, and .ltoreq.4.0-7.0 .mu.mole ethidium bromide to the suspension and increase of the fluorescence intensity is measured in sample with bacteria as compared to the samples without bacteria at 540-620 nm with excitation at 260-320 or 420-530 nm, and the concn. of bacteria is calcd. from the increase of the fluorescence intensity. ICM C12N001-00 ICS C12Q001-00 ICI C12Q001-00, C12R001-01 CC9-5 (Biochemical Methods) Section cross-reference(s): 10 bacteria detn suspension fluorometry ST TT Bacteria Microorganism (detn. of, in suspensions, by fluorometry) IT Spectrochemical analysis (fluorometric, for bacteria in suspensions) IT 1239-45-8 9002-93-1 RL: ANST (Analytical study) (in bacteria detn. in suspension by fluorometry) L17 ANSWER 155 OF 179 CA COPYRIGHT 2003 ACS ИA 103:3115 CA

Identification of bacterial pathogens by laser excited ΤI fluorescence Coburn, J. T.; Lytle, F. E.; Huber, D. M. AU Dep. Chem., Purdue Univ., West Lafayette, IN, 47907, USA CS Analytical Chemistry (1985), 57(8), 1669-73 SO CODEN: ANCHAM; ISSN: 0003-2700 DT Journal English LA The title rapid method relies on the extent of aminopeptidase hydrolysis AB of a series of nonfluorescent L-amino acid .beta.-naphthylamides to produce the highly fluorescent .beta.-naphthylamine by the pathogen of interest. The luminescence background was composed of Raman and Rayleigh scatter, fluorescent impurities in the buffer, .beta.-naphthylamine fluorescence due to substrate decompn., and emission of the biol. matrix. The blank levels were systematically examd. and reduced to levels which allow the measurement of the fluorophore in the 0.1-nM range. Thus, unambiguous identification of pathogens at the 50,000 cell/mL level was achieved. This corresponds to 2-3 orders of magnitude fewer cells than needed by other techniques. Identification of pathogens at this level will drastically reduce the cell growth period from 48 to 6 h and afford a more rapid turnaround time for bacterial identification. 9-5 (Biochemical Methods) CC Section cross-reference(s): 10 bacteria pathogen identification laser fluorometry; ST aminopeptidase profiling bacteria identification Pseudomonas syringae phaseolicola TT Xanthomonas phaseoli (identification of, by aminopeptidase profiling with laser fluorometry) Spectrochemical analysis IT (fluorometric, laser-induced, for aminopeptidase substrates, in bacteria pathogen profiling) ΙT Fluorometers (laser, time-resoln., for bacteria pathogen identification) IT Bacteria (pathogenic, identification of, by aminopeptidase profiling with laser fluorometry) ΙT 716-94-9 720-82-1 729-24-8 732-84-3 732-85-4 740-57-8 3326-64-5 4357-95-3 4420-88-6 7182-70-9 1259-69-4 3326-63-4 14525-44-1 7424-15-9 7424-16-0 16037-15-3 RL: RCT (Reactant); RACT (Reactant or reagent) (hydrolysis of, by aminopeptidase in bacteria pathogen identification by laser fluorometry) 9031-94-1 TΤ RL: ANST (Analytical study) (in bacteria pathogen identification by laser fluorometry) L17 ANSWER 156 OF 179 CA COPYRIGHT 2003 ACS 102:75146 CA ANA comprehensive method for the measurement of fluorescence ΤI lifetime in picoseconds with a phase fluorometer and its application for the determination of excitation enery transfer rate in light-harvesting pigment antenna of green bacteria ΑU Fetisova, Z. G.; Kharchenko, S. G.; Blagoveshchenskii, Yu. N.; Borisov, A. Yu. CS USSR Vestnik Moskovskogo Universiteta, Seriya 16: Biologiya (1984), (4), 56-9 SO CODEN: VMUBDF; ISSN: 0137-0952 DΤ Journal Russian LA AΒ A universal phase fluorometric method for measuring fluorescence lifetime in picoseconds is based on mixing 2 fluorescence beams. The photocathode of the phase fluorometer registers the superimposed 2 light components: the desired short-lived and long-lived components of the same object in a suitable solvent. This method was specially modified for

measuring the fluorescence of photosynthetic pigments (in vivo). The 2-component systems were obtained from 2 solns. of Na fluorescein in 0.001M NaOH, placed in the adjacent cuvettes, 1 of which contained KI as a quencher. Math. equations are given to do the necessary calcns. The method was used to measure the rate of excitation energy transfer in light-harvesting pigment antenna of Chlorobium limicola with a photoreaction center P840. About 90% bacteriochlorophyll c with light-harvesting property participates in transfer of excitation energy transfer to bacteriochlorophyll a and accomplishes it in 20-60 ps with an efficiency of >95% under unsatisfactory light conditions (for photosynthesis). 9-5 (Biochemical Methods) Section cross-reference(s): 10, 11 fluorescence lifetime picosecond detn; energy transfer Chlorobium fluorescence picosecond Energy transfer (fluorescence, in photosynthetic systems of green bacteria, method for measurement of) Pigments, microbial (light-harvesting, excitation energy transfer rate detn. in) Chlorobium limicola (photosynthetic systems of, excitation energy transfer in, method for detn. of) Fluorometry (picosecond, of excitation energy transfer in light-harvesting pigment antenna of green bacteria) Photosynthetic systems (reaction center, P840, of Chlorobium limicola, excitation energy transfer in, method for detn. of) 17499-98-8 53986-51-9 RL: ANST (Analytical study) (of Chlorobium limicola, method for excitation energy transfer rate detn. in relation to) ANSWER 169 OF 179 CA COPYRIGHT 2003 ACS 93:40914 CA Two-parameter analysis of microbial cell constituents Hutter, K. J.; Stoehr, M. Inst. Exp. Pathol., Dtsch. Krebsforschungszent., Heidelberg, 6900, Fed. Rep. Ger. Microbios Letters (1979), 10(39-40), 121-8 CODEN: MILEDM; ISSN: 0307-5494 Journal English Flow cytometry is a new assay to investigate different cellular constituents, e.g., the DNA and protein content of a wide variety of biol. specimens. Investigations have involved quant. fluorescent staining of monodisperse cell populations in liq. suspension at a flow rate of 1000 cells/s. The development of a new dual laser beam excitation device for flow cytometry anal. has enabled simultaneous measurement of the DNA and protein content of baker's yeast cells. The combination of light sources consisted of an Ar ion laser with an excitation wavelength of 488 nm, and a 2nd Ar ion laser at 360 nm wavelength. microbial DNA content was stained by 4',6-diamidino-2-phenylindol while the protein content was fluorochromized by Sulforhodamine 101. staining technique avoided any RNA digestion with pepsin in order to eliminate non-specific cytoplasm fluorescence and revealed a no. of cells in various phases of their life cycle. A correlation between the replicative and metabolic activity was visualized. 9-6 (Biochemical Methods) protein yeast flow cytometry; DNA yeast flow cytometry Saccharomyces cerevisiae (DNA and protein detn. in, by flow cytometry) Deoxyribonucleic acids

. . . .

CC

ST

ΙT

ΙT

IT

ΙT

ΙT

IT

L17 AN

TΙ

ΑU

CS

SO

DT

LA

AB

CC

ST

ΙT

Proteins RL: ANT (Analyte); ANST (Analytical study) (detn. of, in yeast by flow cytometry) 28718-90-3 IT RL: ANST (Analytical study) (DNA in yeast staining with, for detn. by flow cytometry) 60311-02-6 RL: ANST (Analytical study) (protein staining in yeast with, for detn. by flow cytometry) ANSWER 175 OF 179 CA COPYRIGHT 2003 ACS L17 78:56378 CA ANMethod for simultaneous determination of histidine and histamine in ΤI biological liquids. Application to wines Plumas, B.; Sautier, C. ΑU Cent. Rech. Diet., Hop. Bichat, Fr. CS Annales des Falsifications et de l'Expertise Chimique (1972), 65(703), SO 322-36 CODEN: AFECAT; ISSN: 0003-4274 DT Journal LA French Histidine (I) is sepd. from histamine (II) on an Amberlite CG-50 resin; II AB is eluted from the resin with 3N HCl. To det. I, the pH of the effluent is adjusted to 12.15 with NaOH, o-phthalic dialdehyde (C6H4(CHO)2) is added, and after exactly 4 min the condensation reaction is stopped by the addn. of 3N HCl. A fluorescence that is stable for 30 min is obtained; activation 360 m.mu., and emission 450 m.mu.. To det. II, the pH is adjusted to 12.45. The precision and sensitivity of the method for II is 0.05 mg/l. and 0.025 .mu.g/ml, resp., and for I is 0.7 mg/l. and 0.75 .mu.g/ml. Anal. of about 50 wines of different origins gave higher values for I in white wine, 19.43-31.28 mg/l., than for red wines,  $16.99-17.22 \ mg/l$ . White wines had less II,  $2.69-4.90 \ mg/l$ ., than red wines, 6.36-6.49~mg/l. In the course of malolactic reversion, the amt. of I decreased and II increased. This may be due to a simultaneous decarboxylation of I and malic acid. To avoid formation of II, bacteria should be selected that can decarboxylate malic acid without acting on I. 9 refs. CC 16-1 (Fermentations) wine histidine histamine detn; malate wine histamine; fluorometry ST histamine Wine analysis IT(histamine and histidine simultaneous detn. in) TT 71-00-1, analysis RL: ANT (Analyte); ANST (Analytical study) (detn. of, in histamine presence) 51-45-6, analysis TТ RL: ANT (Analyte); ANST (Analytical study) (detn. of, in histidine presence) L17 ANSWER 179 OF 179 CA COPYRIGHT 2003 ACS AN72:107693 CA Direct fluorometric determination of bacterial nucleic acids ΤT Launay, Bernard; Truhaut, Rene ΑU CS Centre Rech. Toxicol., Fac. Pharm., Paris, Fr. Comptes Rendus des Seances de l'Academie des Sciences, Serie D: Sciences SO Naturelles (1969), 269(25), 2614-17 CODEN: CHDDAT; ISSN: 0567-655X DT Journal LAFrench Cultures of Proteus vulgaris and Escherichia coli were collected, washed AB with NaCl 0.16 + citrate 0.01M, and .apprx.20 g wet wt. of each bacterium was suspended in 20 ml of the washing soln. The bacteria were lysed with 0.25-0.5 ml of 20% Na dodecylsulfate in EtOH, and the lysate was diluted 200-fold with Tris-HCl buffer at pH 7.5. Fluorescence

. . .

was detd. in the presence of 8 .mu.g/ml of ethidium bromohydrate with
 excitation at 3400 .ANG. and emission at 5950 .ANG..
 The increase in the fluorescence of ethidium bromohydrate was
 proportional to the vol. of the lysate. The method allowed the detn. of
 DNA at concns. .gtoreq.10-3 .mu.g/ml, corresponding to a level of .apprx.5
 .times. 104 bacteria/ml.
CC 6 (Biochemical Methods)
ST fluorometry nucleic acids; nucleic acids fluorometry
IT Escherichia coli
 (deoxyribonucleic acid detn. in)
IT Nucleic acids, deoxyribo RL: ANT (Analyte); ANST (Analytical study)

(detn. of, in bacteria)

IT Proteus

(vulgaris, deoxyribonucleic acid detn. in)

=>

. . . .